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- (54) NONAQUEOUS ELECTROLYTIC SOLUTION AND LITHIUM SECONDARY BATTERIES
- (57) In order to manufacture a lithium socondary battery having excellent performances in safety under overcharge condition, cycle property, electric capacity, and storage endurance, 0.1 wt.% to 10 wt.% of a tertalky/benzene compound is favorably incorporated into
- a non-aqueous electrolytic solution comprising a nonaqueous solvent and an electrolyte, preferably in comblination with 0.1 wt.% to 1-5 wt.% of a biphenyl compound.

### Description

[Field of Invention]

[0001] The present invention relates to a lithium secondary battery showing high safety under the condition of overcharge, and excellent battery characteristics in cycle property, electric capacity and storage endurance, and further reliates to a non-equeous electrolytic solution favorably employable for preparing the lithium secondary battery.

### [Background of Invention]

[0002] Recently, filtium secondary batteries are generally employed as electric sources for driving small electronic devices. They are also employed as electric sources for driving potable electronic transmission apparatuses such as small size video carriers, ceilsiar phones, and note-size personal computes. They are also expected as electric sources for motor carrs. The filtum secondary battery essentially comprises a positive electrode, a nenequeous electrodic sources for motor carrs. The filtum secondary battery stitizing a positive electrode of a third motor control of the electrodic personal personal positive electrodic personal pers

[0003] In the lithium secondary battery, the positive electrode releases an excessive lithium and the excessive lithium 20 deposits on the negative electrode to produce dendifie, if the battery is overcharged to exceed the ordinary working voltage. Therefore, both of the positive and negative electrodes become unstable. When both electrodes become unstable, the carbonate in the electrolytic solution decomposes upon contact with the unstable electrodes and an exothermic reaction regidy cours. Accordingly, the battery shormally generates thest and safety of the battery fowers. This phenomenon makes increased troublos in the case that the battery generates an electric current of an increased energy density.

[0004] Until now, it has been proposed that an addition of a small amount of an aromatic compound to the electrolytic solution is effective to assure the safety under the overcharge condition.

[0005] JP-Ar-7:002814 describes that an organic compound having a molecular weight of 500 or less and a x-electron orth which gives a reversible oxidation-reduction potential at a potential or note side relative to the posterible electrode openital under the fully charged condition, which is represented by an anisole derivative, is used as an additive for an electrividic activities obtain.

[0006] JPA-2000-158233 describes that an organic compound having a x-electron onto which gives a reversible softential to the positive electron potential at a potential of noble side relative to the positive electrode potential under the fully charged condition, which is represented by an anisole derivative, biphenyl, end, 4.4-denethy biphenyl, is used as an additive for an electrohic solution. It is described that the organic compound such as the above-mentioned anisole derivative or biphenyl derivative generates a redox shuttle in the battery, so that the safety of the battery is assured. [0007] JPA-6-106835 (corresponding to United States Patents, 578, 934) describes a measure for assuring sefety of a battery under the overcharge condition by increasing an internal resistance in the battery. The increase of an internal resistance can be accordinated under the promoters and the promoters are supported to the promoters of the

40 furan in an amount of approx. 1 to 4% to polymorize the compound such as biphenyl at a voltage exceeding the maximum working voltage of the battery.

[0008] JP-A-9-171940 (corresponding to United State Patents 5,776,827 and 5,033,797) also describes a measure for assuring safety of a battery under the overcharge condition, by working an internal current breaker has the battery. The linternal current breaker can be worked using a monomer such as biphenyl, 3-R-thiophene, 3-chiorothiophene or furan in an amount of approx. 1 to 4% to polymerize a compound such as biphenyl and produce a gas at a voltage exceeding the maximum working voltage of the battery.

[0009] JP-A-10-221258 also describes a measure for assuring safety of a battery under the overcharge condition, by producing an electroconductive polymer in the battery. The production of an electroconductive polymer and be accomplished by using a monomer such as biphenyl, 3-R-thiophene, 3-chlorothiophene or furan in an amount of approx. I to 4% to polymerize the compround at a votage occeeding the maximum working voltage of the battery.

[0010] IPA-10-275632 describes that a noninor aromatic compound having on ally or the obstery. [0010] IPA-10-275632 describes that a noninor aromatic compound having an alkyl group is incorporated into an organic electrolytic solution of a secondary battery which comprises a linear ester as a main solvent. As the nonionic aromatic compound having an alkyl group, there are mentioned a trimelitic ester, tri-2-ethylh trimelitiate, dimethyl phthalate, dibutyl phthalate, butylbenzene (normal, terilary, or iso), cyclohexylbenzene and toluene.

5011] JPA-11-16251 (corresponding to United States Patent 6,074,777) describes that the addition of biphenyl or the like is a pict to lower the battery performances used has cycler properly when the battery is subjected to respected cyclic. The procedure in which the battery is charged to a voltage exceeding the maximum voltage of 4.1 V, or the battery is charged and discharged at a high temperature such as 40°C or higher for a join poreirod of time, and that heese recibients are recibients are:

more apparently noted when the additive is added in an increased amount. This publication further describes that an electrophytic solution combaining 2,2-diphenytipropane or its analogous compound is teverably employed for assuring the selfey of a battery under the overlarge condition because 2,2-diphenytipropane or its analogous compound polymerizes to generate a gas, resutting in working of an internal current breaker, or to give an electroconductive polymer, resulting in generation of internal short-directly.

[0012] The anisole derivatives and biphenryl derivatives described in JP-A-7-02514 and JP-A-2000-156243 favorably work under the condition of overcharge by utilizing redox shuttle, but give adverse effects to the cycle property and storage endurance. In more detail, the anisole derivatives and obliphenryl derivatives gradually decompose when the battery is subjected to the repeated charge-discharge procedure, if they are locally subjected to an elevated voltage in the case that the battery is used at a lamperature of 40°C or higher, or that the buttlery is used at an ordinary working voltage. Therefore, the contents of the anisole derivative and biphenryl derivatives decrease by decomposition in the case that end of south and the desired safety cannot be assured when the charge-discharge procedure is certified us the end of the desired and ordinary decreases and composition of the carried out ether 200 cycle charge-discharge procedure is expected.

[0013] Likewise, bithemyl, 3-h-hischene, 3-chlorothiophene, and furan which are described in JP-A-9-108835, JP-A-9-171640, and JP-A-11-62512, livey live adverse effects to the cycle properly and storage endurance. These problems are more preminently noted when the amount of biphenyl increases. In more detail, since biphenyl or the file decomposes by oxidation at a potential of 4.5 or lower, the content of biphenyl or the file gradually decreases when it is focally subjected to somewhat high voltage in the course of working at 40°C or higher or at an ordinary working voltage, resulting in decrease of the cycle iffe. Further, since the content of biphenyl or the file gradually to decomposition, the desired safely is sometimen not assured when the charge-discharge procedure is carried out fairs the 300 cycle charge-discharge procadue is repeated.

[0014] In addition, a battery containing 22-diphenylpropens and its analogous compound (which is described in JPA-11-1625); shows only unsatisfactory safety under the overcharge condition, but the strained safety is highly than a battery having no such additive. On the other hand, it is known that the battery oritaining 22-diphenylpropase and its analogous compound shows a cycle properly better than a battery oritaining piphenyl, but worse than a battery containing no additive. Thus, if a battery showing a cycle properly better than that aims duy the battery containing piphenyl is required, safety should be sacrificed.

[0015] It is an object of the present invention to provide a lithium secondary battary showing high safety under the oversharge condition, and excelent battery performances in cycle property, electric capacity and storage endurance, and turther provide a non-equeous electrolytic solution fevorably employable for preparing the lithium secondary battery.

### [Disclosure of Invention]

[0016] The invention resides in a non-equeous electrolytic solution for lithium secondary betteries comprising a nonequeous solvent and an electrolyte, which further contains 0.1 wt% to 10 wt.% of a tert-alky/benzene compound and 0.1 wt.% to 1,5 wt.% of a biphenyl compound.

[01:8] The biphenyl compound to be incorporated into the non-aqueous electrolytic solution is preferably represented by ε<sup>2</sup>-4<sup>3</sup> in which each of ε<sup>3</sup> and ε<sup>3</sup> is a barzene ring that may have 1 to 5 substituents on the ring. Examples of the biphenyl compounds include biphenyl, d-ariphenyl, n-terphenyl p-temphenyl, 4-methylbiphenyl, 4-methylbipheny

50 [0019] Further, the invention resides in a non-aqueous electrolytic solution for lithium secondary batteries comprising a non-aqueous solvent and an electrolyte, which further contains 0.1 nt. % to 20 nt. % of a tert-airly/spenzee compound (in which the tert-airly/spenze) as to 13 catton anoms). A representative example of the tert-airly/spenzee compound is left-perty/lenzeen. In other words, the tert-airly/spenzeen compound having 5 to 13 carbon atoms can be employed for accomplishing the object of the invention which the biblewhyl compound.

[0020] Furthermore, the invention resides in a lithium secondary battery comprising a positive electrode comprising a compound oxide containing lithium and coals, linkel or manganese, a negative electrode comprising lithium material, lithium aloy or material capable of roceiving and releasing lithium, and a non expectous electrolytic solution of the above-mentioned invention.

[0021] As described hereinbefore, the previously known methods for obviating overcharge are as follows:

a method of conducting a redox shuttle at a voltage of approx. 4.5 V;

a method of increasing an internal resistance of a battery by polymerizing an additive at a voltage of 4.5 V or lower; a method of forming short-circuit by generating a gas to work an internal current breaker; and a method of assuring safety of a battery under the overcharge condition by producing a polymer to generate internal

short-circuit.

[0022] In contrast, the present invention utilizes a non-aqueous electrolytic solution containing a tert-alkylbenzene compound as an additive. The mechanism of the invention for assuring safety under the overcharge condition is considored as follows: the tort-alky/benzene compound decomposes by oxidation at a potential of +4.6 V to +5.0 V (relative value to that of lithium), and cobalt or nickel in the positive electrode rapidly dissolves and deposits on the negative electrode to inhibit a reaction of a carbonate in the non-aqueous electrolytic solution with a lithium metal deposited on the negative electrode.

[0023] Further, in the invention, the internal short circuit may be formed in the battery by the deposition of cobalt or nickel, whereby the overcharge inhibitive effect can be attained and the safety of battery can be assured.

[9024] Furthermore, the addition of a small amount (0.1 wt.% to 1.5 wt.%) of a biphenyl compound together with the tert-alkylbenzene compound increases the overcharge inhibitive effect of the tert-alkylbenzene compound. The addition of a small amount of a biphenyl compound surprisingly further improve the battery performances which has not been expected from the previously accepted knowledge.

100251 Moreover, since the tert-alkylbenzene compound contained in the non-aqueous electrolytic solution has such a high exidation potential as +4.6 V to +5.0 V (relative value to the exidation potential of lithium), the tert-alky/benzene compound does not decompose under a locally occurring high voltage such as higher than 4.2 V in the case that the battery is used at such a high temperature as 40°C or higher, and the charge discharge procedure is repeated at an ordinary voltage.

[0026] The addition of a small amount (0.1 wt.% to 1.5 wt -%) of a biphenyl compound alone cannot inhibit overcharge of a battery, it is discovered, however, that it is effective to improve the battery performances, when it is employed in combination with a tert-alkylbenzene compound. This is considered that the decomposition of the biphenyl compound is low. Further, since the tert-alkylbenzene brings about an effect to inhibit overcharge, the safety of a battery is ac-

complished even in the case that a 300 cycle overcharge test is carried out. Therefore, it is considered that a lithium secondary battery showing not only the safety under the overcharge condition but also high battery performances in the cycle property, electric capacity and storage endurance can be manufactured.

# [Detailed Description of Invention]

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[0027] Examples of the tert-alkylbenzene compounds to be dissolved in an electrolytic solution comprising a nenaqueous solution and an electrolyte include the following compounds:

tert-buty/benzene, 1-fluoro-4-tert-buty/benzene, 1-chloro-4-tert-buty/benzene, 1-bromo-4-tert-buty/benzene, 1-lodo-4-tert-butylbonzone, 5-tert-butyl-m-xylene, 4-tert-butyltoluene, 3,5-di-tert-butyltoluene, 1,3-di-tert-butylbonzene, 1,4-di-tert-buty/benzene, 1,3-5-tri-tert-buty/benzene, tert-penty/benzene, (1-ethyl-1-methylpropyl) benzene, (1,1-diethylpropyl)benzene, (1,1-dimethylbutyl)benzene, (1-ethyl-1-methylbutyl) benzene, (1-ethyl-1-ethylbutyl) benzene, (1,1,2-trimethylpropyl)benzene, 1-fluoro-4-tert-pentylbenzene, 1-chloro-4-tert-pentylbenzene, 1-bromo-4-tert-pentylbenzene, 1-iodo-4-tert-pentylbenzene, 5-tert-pentyl-m-xylene, 1-methyl-4-tert-pentylbenzene, 3,5-ditert-pentylloluene, 1,3-di-tert-pentylbenzene, 1,4-di-tert-pentylbenzene, and 1,3,5-tri-tert-pentylbenzene.

[0028] The tert-alkylbenzene compounds can be employed singly or in combination of two or more compounds. [0029] A preferred tert-alkylbenzene compound to be added to the non-aqueous electrolytic solution of the invention

is a compound having the formula of (R1) (R2) (R3)Co1 [in which R1 is an alkyl group of 2 to 4 carbon atoms, each of R2 and R3 Independently is an alkyl group of 1 to 4 carbon atoms, and §1 represents a benzene ring that may have 1 to 5 substituents on the ring). The use of this compound is effective to improve particularly the cycle property of the non-aqueous electrolytic solution.

[0030] In the formula of (R1) (R2) (R3)C-\$\phi\_1\$, R1 preferably is an alkyl group such as ethyl, propyl or butyl, and each of F2 and R3 independently is an alkyl group such as methyl, ethyl, propyl, or butyl. The alkyl group can be a linear or branched alkyl group.

[0031] In the benzene ring that may have 1 to 5 substituents on the ring which is represented by \$\phi^1\$, each of the substituents preferably is independently a linear alkyl group such as methyl, ethyl, propyl or butyl, or a branched alkyl group such as isopropyl, isobutyl, sec-but tert-butyl, or tert-pentyl. Otherwise, the substituent can be a cycloalkyl group

- having 3 to 6 carbon atoms such as cyclopropyl or cyclohesyl; phenyl; benzyl; or an alkylated phenyl or benzyl group such as toly, Inch-butyphenyl, tert buybbenzyl, or test pentylphenyl; a halogen atom such as fluorine, chlorine, bromine, or iodine. The benzene ring prelentidy has a bydrocarbly group having 1 to 12 carbon atoms or a halogen atom.
- [0332] Examples of the above-mentioned far-fally/tenzane compounds include tert-penty/benzene, (1-ditty/-1-methy/propy/)benzene, (1-ditty/-propy/)benzene, (1-ditty/-ybroy)/benzene, (1-ditty/-ybroy)/benzene, (1-ditty/-ybroy)/benzene, (1-ditty/-ybroy)/benzene, (1-ditty/-ybroy)/benzene, (1-ditty/-ybroy)/benzene, (1-ditty/-distry-penty/benzene, and (1-1,2-dittenty/-ybroy)/benzene, (1-ditty/-distry-penty/-ybroy)/benzene, (3-ditty-penty/-ybroy)/benzene, (1-ditty-penty/-benzene, 1-ditty-distry-penty/-benzene, 1-ditty-distry-penty/-benzene, 1-ditty-distry-penty/-benzene, 1-ditty-distry-penty/-benzene, 1-ditty-distry-penty/-benzene, 1-ditty-distry-penty/-benzene, 1-ditty-distry-penty/-benzene, 1-ditty-penty/-benzene, 1-ditty-
- [0033] Exemples of the biphenyl compounds include biphenyl, o-terphenyl, m-terphenyl, p-terphenyl, 4-methylbiphenyl, 4-cthylbiphenyl, and 4-tert-butylbiphenyl.
  - [0034] Particularly, if a portion of the tert-buty/benzene or the like having a high exidation potential of 4.8 to 6.0 V is replaced with a biphenyl compound having a low exidation potential of 4.5 V (e.g., o-terphenyl), the safety under the overcharce condition is enhanced.
- 15 [0035] If a portion of the tert-alkythenzene compound is replaced with the biphenyl compound, the content of the tert-alkythenzene preferably is 10 times or lower, more preferably 0,3 to 5 times, particularly preferably 0,5 to 3 times, as much as the content of the biphenyl compound.
- [0036] As described hereinbefore, the combination of the tert-allythenzene compound and biphenyl compound which have different oxidation potential is effective to improve the security under the overcharge condition and the battery performances.
  - [0037] If the amount of the lart-skyldenzene compound is excessive, the electrolytic solution may show a different electroconductivity and the battery performances may lower if the sensors is excessively low, enough salety-improving effect cannot be accomplished. Accordingly, the amount preferably is in the range of 0.1 to 10 wtr.5, more preferably 16.6 btt.5, besed on the amount of the electrolytic coultion.
- 25 [0038] If the amount of the biphenyl compound is excessive, the biphenyl may decompose in the battery even under ordinary working conditions, and the battery performances may lower. If the amount is excessively low, enough safety-improving effect, and battery performances cannot be accomplished. Accordingly, the amount preleably is in the amount of the electrolytic solution.
- [0039] Examples of the non-equeous solvents employable for the preparation of the non-equeous electrolytic solution of the invention include cyclic carbonates such as ethylene carbonate (EC), propylene carbonate (PC), but/yiere carbonate (PC), and vinylene carbonates (PC) is a directly carbonate (PC), and vinylene carbonates where a special carbonates (PC), and vinylene carbonates (PC), and distript carbonate (PC); ethers such as tetrahydrofuran, 2-methyltrathyldrofuran, 14-directly carbonate (PC); ethers such as retrahydrofuran, 2-directly carbonate (PC); ethers such as acetonitrile; esters such as mathyl propionate, methyl phratiale, and corty phratiale, and cardiose such as directly directly carbonates and 12-directly propionate, methyl phratiale, and corty phratiale, and cardiose such as directly tomative the propionate, methyl phratiale, and corty phratiale, and cardiose such as directly tomatics.
  - [0040] The non-equeous solvents can be employed singly or in combination of two or more solvents. There are no imitiations with respect to possible combinations. Examples are combinations of a cyclic carbonate and a fine
- 40411 Examples of the electrolytes to be dissolved in the non-equeous solvent for the preparation of a non-equieous electrolyte solution hoticle LiPE, LIBE, LIPE, L
- [0042] The electrolytic solution can be prepared, for instance, by mixing the above-mentioned non-aqueous solvents; dissolving the electrolytic in the mixture; and further dissolving at least one tert-alkyticenzene compound and optionally at least one bipheryl compound in the mixture solution.
  - [0043] The electrolytic solution of the Invention is favorably employable for manufacture of a secondary battery, particularly lithium secondary battery. There are no limitations with respect to materials of the secondary battery other than the electrolytic solution, and various known materials can be employed.
- Odd4] The positive electrode active material preferably comprises a compound metal coids containing litikism and cobalt or nicket. Examples of the compound metal coidse include LEGO<sub>2</sub>, LNNO<sub>2</sub>, COI CH C+L), and LIMn<sub>2</sub>O<sub>4</sub>, Fruither, an appropriate initiative such as a miniture of LEGO<sub>2</sub> and LIMn<sub>2</sub>O<sub>4</sub>, a miniture of LEGO<sub>2</sub> and LINn<sub>2</sub>O<sub>4</sub>, and thing of LEGO<sub>2</sub> and LINn<sub>2</sub>O<sub>4</sub> and thing of LEGO<sub>2</sub> and LINn<sub>2</sub>O<sub>4</sub>.
- [0045] The positive electrode can be manufactured by kneading the above mentioned positive electrode active massistatia, an electro-conductive material such as acatylane black or carbon black, and a binder such as polyticiralluor-cethylene) (PTFE), polytymickiene fluoride) (PVFF), styrene-busidiene copolymer (SBR), acryloniri.bintidadee ocpolymer (NBR) or carboxymethycetiluloses (CMC) to give a positive electrode composition; coaling this positive electrode composition on a collector such as aluminum foli, stainless foli, or faith plate; drying the coarted composition; pressing

the dried composition; and heating the pressed composition under vacuum at a temperature of approximately 50 to 2500C for approximately 2 hours.

IDA6] As the negative electrode (negative electrode active material), altihlum metal, a lithium alloy, a centronascous, material which, can also the and release lithium (e.g., thematy decomposed carbonascous material, oclor, can also that of release lithium (e.g., thematy decomposed carbonascous materials) expenses a stifficial graphite and natural graphile, fired organic polymer, and carbon fiberl, or a compound the oxide can be employed. It is preferred to employ a carbonascous material harding a graphite organic statuture in which the lattice distance of lattice surface (002), namely, deg., Is in the range of 0.335 to 0.340 mm (nanometer). The negative electrode active material in the providery forms such as carbonascous powder is preferrably used in combination with a brider such as ethic material in the providery client combination with a brider such as ethic propriets of the combination of the brider such as ethic propriets of the combination of the provider of the combination of the combination of the provider of the combination of

[0047] There are no appecific limitations with respect to the structure of the lithium secondary battery of the invention. For instance, the secondary, battery can be a battery of coint hype competing a positive electrode, a negative electrode and a separator in a single or ptural layers, or a cylindrical or prismatic battery comprising a positive electrode, a negative electrode, and as expandor roll. The separator can be made of microporous film, wovan fabric or non-woven fabric.

[0448] The lithium secondary battery of the invention shows good cycle property for a long period of time even if it is used to work at a maximum working vallage of 4.2 or higher, and the good cycle property can be maintained even if it is used to work at an eximum working vallage of 4.3 V. The cut-off voltage can be set to 2.0 V or higher, the can be set to 2.6 V or higher, there is no limitation with respect to a current value, but a constant current of 0.1 to 9.0 to generally utilized. The battery of the invention can be charged and discharged in such a wide temperature range of 4.0°C to 100°C, However, the temperature prefereably the within to 10 gain.

### (Example 1)

25 1) Preparation of electrolytic solution

[0049] In a non-equeous mixture of EC/PC/DEC=30/5/65 (volume ratio) was dissolved LIPF<sub>6</sub> to give an electrolytic solution of 1M concentration. Subsequently, 2.5 wt.% of tert-buty/ibenzene and 0.9 wt.% of biphenyl were added to the electrolytic solution.

2) Manufacture lithium secondary battery and measurement of its battery performances

[0050] LiCoO<sub>2</sub> (positive electrode active material, 90 vs.%), acetylene black (electro-conductive material, 5 vs.%), and poly(vinylidene fluoride) (binder, 5 vst.%) were mixed. To the resulting mixture further added 1-metryl-2-pyrrolidone, to drive a sturry. Thus produced sturry was coaled on alturnitum foil, dried, and pressed to give a positive electrode.

[0051] Artificial graphite (negative electrode active material, 95 wt. %) and poly (virylidene fluoride) (binder, 5 wt. %) were mixed. To the resulting mixture was further added 1-methyl-2-pyrolidione to give a sturry. Thus produced sturry was coated on copper fol. died, and pressed to give a neutrive electrode.

[0052] A microporous polypropylene film separator and the above-mentioned electrolytic solution were placed in a cylindrical case to give a cylinder-type battery of 18650 size (diameter: 18 mm, height: 65 mm). To the battery were provided a pressure reloasing port and an internal current breaker.

[0033] For performing a cycle text using the 18650 battery, the battery was charged at an elevated temperature (45°C) with a constant electric current of 1.45 A, 10°C) under a constant electric constant electric current of 1.45 A, 10°C) under a constant electric current 1.45 A (10°C) to give a terminal voltage of 2.5 V. The charm-discharge cells was repeated.

[0054] The initial discharge capacity was almost the same as the capacity measured in a battery using an 1M LiPF<sub>8</sub> and EC/PC/DEC (30/5/65, volume ratio) solvent mixture [Comparison Example 1].

[0055] After a 300 cycle charge-discharge procedure, the retention of discharge capacity was 84.4% of the initial discharge capacity (100%). Further, the battery showed good high temperature storage endurance.

[0066] Subsequenty, the 18650 battey having been subjected to the 300 charge-discharge cycle test was excessively charged at a room temperature (20°C) from a fully charged condition using a constitut current of 2.9 A (2C), for evalualing a resistance to the overcharging, it was confirmed that the current broke after 22 minutes, and the highest surface temperature of the battery after the breakage of current was 6°TC.

[0057] The materials of the tested 18560 size cylinder battery and the battery performances are set forth in Table 1.

### [Example 2]

[0058] The procedures of Example 1 were repeated except that the amount of biphenyl in the electrolytic solution

was changed to 0.5 wt.%, to prepare a cylinder battery.

[0059] The materials of the tested battery, the discharge capacity retention ratio after the 300 cycle charge discharge procedure, a period of time until breakage of current occurs, and the highest surface temperature of the battery after the breakage of current are shown in Table 1.

#### (Example 3)

[0060] The procedures of Example 1 were repeated except that the amount of biphenyl in the electrolytic solution was changed to 1.3 wt.%. to prepare a cylinder battery.

[0051] The materials of the tested battery, the discharge capacity retention ratio after the 900 cycle charge-discharge procedure, a ported of time until breakage of current occurs, and the highest surface temperature of the battery after the breakage of current are shown in Table 1.

### [Example 4]

[0062] The procedures of Example 1 were repeated except that the amount of biphenyl in the electrolytic solution was replaced with 0.9 wt.% of o-terphanyl, to prepare a cylinder battery.

[0063] The materials of the tested battery, the discharge capacity retention ratio after the 300 cycle charge-discharge procedure, a period of time until breakage of current occurs, and the highest surface temperature of the battery after the breakage of current are shown in Table 1.

### [Example 5]

[0064] The procedures of Example 1 were repeated except that the text-butylbenzene was replaced with 2.5 wt.% of text-pentylbenzene and the hiphenyl was replaced with 0.9 wt.% of 4-ethylbiphenyl in the electrolytic solution, to prepare a cylinder battery.

[0065] The materials of the tested battery, the discharge capacity retention ratio after the 300 cycle charge-discharge procedure, a period of time until breakage of current occurs, and the highest surface temperature of the battery after the breakage of current are shown in Table 1.

### [Example 6]

[0066] The procedures of Example 1 were repeated except that tert-burybenzene and tert-pentylbenzene were employed as tert-alkylbenzene compounds in an amount of 2 wt. % for each, and 4-metrybliphenry was employed as the biblientyl compound in an amount of 0.5 wt.5 in the electrolytic solution, to prepare a cylinder batch.

[0067] The materials of the tested battery, the discharge capacity retention ratio after the 300 cycle charge-discharge procedure, a period of time until because of current occurs, and the highest surface temperature of the battery after the breakage of current are shown in Table 1.

### 40 [Comparison Example 1]

[0068] The procedures of Example 1 were repeated except that neither tert-alky/benzene compound nor biphenyl compound was incorporated, to prepare a cylinder battery.

[0089] The materials of the tested battery, the discharge capacity retention ratio after the 300 cycle charge-discharge procedure, a period of time until breakage of current occurs, and the highest surface temperature of the battery after the breakage of current are shown in Table 1.

# [Comparison Example 2]

90 [0070] The procedures of Example 1 were repXpeated except that biphenyl was employed in an amount of 1.3 wt. % in the electrolytic solution and no tert-ally/benzene compound was incorporated, to prepare a cylinder battery, [0071] The materials of the tested battery, the discharge depactly retention ratio after the 300 cycles charge-discharge.

procedure, a period of time until breakage of current occurs, and the highest surface temperature of the battery after the breakage of current are shown in Table 1.

### [Comparison Example 3]

[0072] The procedures of Comparison Example 2 were repeated except that biphenyl was employed in an amount

of 4 wt.% in the disctripylic solution and no tert-ally/benzene compound was incorporated, to prepare a cylinder battery, [073] The materials of the tested battery the discharge capacity retention in also after the 300 cycle charge-discharge procedure, a period of time until breakage of cornect occurs, and the highest surface temperature of the battery after the breakage of current are shown in Table 4

### [Example 7]

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- [0074] The procedures of Example 5 were repeated except that LICoO<sub>2</sub> (positive electrode active material) was replaced with LiNi<sub>0.6</sub>Co<sub>0.2</sub>O<sub>2</sub>, to prepare a cylinder battery.
- [0075] The materials of the tested battery, the discharge capacity retention ratio after the 300 cycle charge-discharge procedure, a period of time until breakage of current occurs, and the highest surface temperature of the battery after the breakage of current are shown in Table 1.

### (Comparison Example 4)

[0076] The procedures of Example 7 were repeated except that neither test-alkybenzene compound nor biphenyl compound was incorporated, to prepare a cylinder battery. The battery performances were evaluated. [0077] The materiaks of the tested battery and the battery performances are set forth in Table 1.

### 20 [Example 8]

- [0078] The procedures of Example 1 were repeated except that the tert-butylbenzene was replaced with 4-fluoro-tert-pentylbenzene in an amount of 3.0 wt.% in the electrolytic solution, to prepare a cylinder battery.
- [0079] The materials of the tested battery, the discharge capacity retention ratio after the 300 cycle charge-discharge procedure, a period of time until breakage of current occurs, and the highest surface temperature of the battery after the breakage of current are shown in Table 1.

## [Comparison Example 5]

- [080] The procedures of Comparison Example 1 were repeated except that 3.0 wt.% of toluene and 0.5 wt.% of biphenyl were employed in the electrolytic solution, to prepare a cylinder battery.
  - [0081] The materials of the tested battery, the discharge capacity retention ratio after the 300 cycle charge-discharge procedure, a period of time until breakage of current occurs, and the highest surface temperature of the battery after the breakage of current are shown in Table 1.

#### [Comparison Example 6]

- [0082] The procedures of Comparison Example 1 were repeated except that 3.0 wt.% of n-butylbenzene and 0.5 wt. % of biphenyl were employed in the electrolytic solution, to prepare a cylinder battery.
- [0083] The materials of the tested battery, the discharge capacity retention ratio after the 300 cycle charge-discharge procedure, a period of time until breakage of current occure, and the highest surface temperature of the battery after the breakage of current are shown in Table 1.

### [Comparison Example 7]

- [0084] The procedures of Comparison Example 1 were repeated except that 3.0 wt.% of di-n-butyl phthalate and 0.5 wt.% of biphenyl were employed in the electrolytic solution, to prepare a cylinder battery.
- [0085] The materials of the tested battery, the discharge capacity retention ratio after the 300 cycle charge-discharge procedure, a period of time until breakage of current occurs, and like highest surface temperature of the battery after the breakage of current are shown in Table.

# [Comparison Example 8]

- [0086] The procedures of Comparison Exemple 1 were repeated except that 3.0 wt.% of 4-fluorotoluene and 0.5 wt. % of biphanyl were employed in the electrolytic solution, to prepare a cylinder battery.
- [0087] The materials of the tested battery, the discharge capacity retention ratio after the 300 cycle charge-discharge procedure, a period of time until breakage of current occurs, and the highest surface temperature of the battery after the breakage of current are shown in Table 1.

# Table 1

5		Positive electrode Negative electrode	Tert-alkyl-benzene (wt.%) Biphenyl(wt. %)	Electrolytic solution (vol.ratio)	Current breakage High, temperature Discharge capacity retention
	Ex.1	LICoO <sub>2</sub>	tert-butyl- benzene (2.5)	1M LiPF <sub>6</sub>	22 min.
10		Graphite		EC/PC/DEC	67°C
			biphenyl (0.9)	(30/5/65)	84.4%
	Ex.2	LiCoO <sub>2</sub>	tert-butyl- benzene (2.5)	1M LiPF <sub>6</sub>	24 min.
15		Graphite		EC/PC/DEC	69°C
			biphenyf (0.5)	(30/5/65)	84.0%
20	Ex.3	LiCoO <sub>2</sub>	tert-butyl-benzene (2.5)	1M LIPF <sub>6</sub>	20 min.
		Graphite		EC/PC/DEC	66°C
			biphenyl (1.3)	(30/5/65)	82.7%
25	Ex.4	LICoO <sub>2</sub>	tent-butyl-benzene (2.5)	1M LIPF <sub>6</sub>	22 min.
		Graphite		EC/PC/DEC	67°C
			o-terphenyl (0.9)	(30/5/65)	84.3%
30	Ex.6	LiCoO <sub>2</sub>	tert-pentyl-benzene (2.5)	1M LIPF <sub>6</sub>	21 min.
		Graphite		EC/PC/DEC	66°C
			4-ethyl biphenyl (0.9)	(30/5/65)	82.3%
35	Ex.6	LiCoO <sub>2</sub>	tert-butyl-benzene (2)	1M LIPF <sub>6</sub>	21 min.
		Graphite		EC/PC/DEC	66°C
			tert-pentyl-benzene (2)	(30/5/65)	85.1%
40			4-methyl-biphenyl (0.5)		
	Com. Ex.1	LiCoO <sub>2</sub>	None	1M LIPF <sub>6</sub>	31 min.
		Graphite		EC/PC/DEC (30/5/65)	thermal runaway 82,8%
45	Com. Ex.2	LICoO <sub>2</sub>	biphenyl	1M LiPF <sub>6</sub>	31 min.
		Graphite	(1.3)	EC/PC/DEC (30/5/65)	thermal runaway 78.3%
	Com. Ex.3	LiCoO <sub>2</sub>	biphenyl	1M LiPF <sub>6</sub>	18 min.
50		Graphite	(4)	EC/PC/DEC (30/5/65)	83°C 72.1%
	Ex.7	LiNi <sub>0.8</sub>	tert-pentyl-benzene (2.5)	1M LiPF <sub>8</sub>	21 min.
		CO <sub>0.2</sub> O <sub>2</sub>		EC/PC/DEC	67°C
55		Graphite	4-ethyl biphenyl (0.9)	(30/5/65)	82.5%

Table 1 (continued)

	tuble 1 (continued)					
	Positive electrode Negative electrode	Tert-alkyl-benzene (wt.%) Biphenyl(wt %)	Electrolytic solution (vol ratio)	Current breakage High temperature Discharge capacity retention		
Com. Ex.4	LiNi <sub>0,8</sub>	None	1M LIPF <sub>6</sub>	31 min.		
	CO <sup>0 2</sup> O <sup>2</sup>		EC/PC/DEC	thermal runaway		
	Graphite		(30/5/65)	80.4%		
Ex.8	LiCoO <sub>2</sub>	4-fluoro-tert- pentylbenzene (3)	1M LIPF <sub>6</sub>	23 min.		
	Graphite	biphenyl (0.5)	EC/PC/DEC	58°C 84.3%		
			(30/5/65)			
Com. Ex.5	LiCoO <sub>2</sub>	toluene (3)	1M LIPF <sub>6</sub>	31 min.		
	Graphite	biphenyl (0.5)	EC/PC/DEC	thermal runaway 81.2%		
			(30/5/65)			
Com. Ex.6	LiCoO <sub>2</sub>	n-butyl-benzene (3)	1M LiPF <sub>6</sub>	31 min.		
	Graphite	biphenyt (0.5)	EC/PC/DEC	thermal runaway 80.1%		
			(30/5/65)			
Com. Ex.7	LICoO <sub>2</sub>	di-n-butyl-phthalate (3)	1M LIPF <sub>6</sub>	31 min.		
	Graphite	blphenyl (0.5)	EC/PC/DEC	thermal runaway 78.4%		
			(30/5/65)	•		
Com. Ex.8	LiCoO <sub>2</sub>	4-fluoro-toluene (3)	1M LIPF6	25 min.		
	Graphite	biphenyl (0.5)	EC/PC/DEC	thermal runaway 79.8%		
			(30/5/65)			
Remark	Remarks: Current breakage: period of time to break current					
	High, temperature: Highest temperature of battery					
	Discharge capacity retention: ratio of discharge capacity after 300 cycle					
Graphit	Graphite: Artificial graphite					

40 [0086] In Examples 1 to 5, a sufficient amount of cobait or nickel deposited on the negative electrode under the overcharge condition. It is clear that a lithium secondary batlery utilizing a non-aqueous electrolytic solution which contains a tert-alkylbenzene compound and a biphenyl compound according to the invention shows safety under the overcharge condition and the cycle property better than the secondary batteries of Comparison Examples.

### 45 [Example 11]

- 1) Preparation of electrolytic solution
- [0089] In a non-squeous mixture of EC/PC/DEC=30/5/65 (volume ratio) was dissolved LIPF<sub>8</sub> to give an electrolytic solution of 1M concentration. Subsequently, 2.0 wt.% of tert-porty/bonzene was added to the electrolytic solution.
  - 2) Manufacture lithium secondary battery and measurement of its battery performances

[0999] LCoO<sub>2</sub> (positive electrode active material, 80 at %), acetylene black (electro-conductive material, 10 at %), and polyfortyldene fluoride) [blinder, 10 at %) were mixed. To the resulting mixture further added 1-methyl-2-gyrnolidone, to give a slurry. Thus produced slurry was coaled on aluminum foli, dried, and pressed to give a positive electrode.

- [0091] Artificial graphite (negative electrode active material, 90 art.%) and poly (vinylidene fluoride) (binder, 10 wt. %) were mixed. To the resulting mixture was further added 1-methyl-2-pyrrolidene to give a sturry. Thus produced slurry was coated on cooper fold, died and pressed to give a negative dectrode.
- [0092] A microporous polypropylene film separator and the above-mentioned electrolytic solution were placed in a coin case to give a coin-type battery (diameter: 20 mm, thickness: 3.2 mm).
- [0093] The coin battery was churged at a room temperature (20°C) with a constant electric current of 0.8 mA under a constant voltage to reach 4.2 V for 5 hours. Subsequently, the battery was discharged to give a constant electric current 0.8 mA to give a terminal voltage of 2.7. The charge-click-parge cycle was repeated.
- [0094] The Initial discharge capacity was almost the same as the capacity measured in a battery using an 1M LIPF<sub>6</sub> on and EC/PC/DEC (30/5/65, volume ratio) solvent mixture containing no tert-alkylbenzene compound [Comparison Example 1].
  - [0095] After a 50 cycle charge-discharge procedure, the retention of discharge capacity was 92.8% of the initial discharge capacity (100%). Further, the battery showed good low temperature storage endurance.
  - [0096] The materials of the coin battery and the battery performances are set forth in Table 2.

(Example 12)

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[0097] The procedures of Example 11 were repeated except that the amount of tert-penty/benzonc in the non-aqueous electrolytic solution was changed to 5.0 wt.%, to prepare a coin battery.

[0098] After the 50 cycle charge-discharge procedure, the retention of discharge capacity was 91.5%.

[0099] The materials of the coin battery and the battery performances are set forth in Table 2.

[Example 13]

- 25 [0100] The procedures of Example 11 were repeated except that the amount of tert-pentylbenzene in the non-aqueous electrolytic solution was changed to 0.5 wt.%, to prepare a coin battery.
  - [0101] After the 50 cycle charge-discharge procedure, the retention of discharge capacity was 90.3%.
  - [0102] The materials of the coin battery and the battery performances are set forth in Table 2.

30 [Comparison Example 11]

- [0103] In a non-aqueous mixture of EC/PC/DEC-30/6/65 (voltume ratio) was dissolved LIPF<sub>6</sub> to give a non-aqueous electrolytic solution of 1M concentration. In this procedure, no ter-distybezene compound was added. [0104] A coin battery was manufactured in the same manner as in Exemple 11, except for employing the above-
- obtained non-aqueous electrolytic solution, and subjected to the evaluation of battery performances.

  [0105] After the 50 cycle charge-discharge procedure, the relention of discharge capacity was 82,6%.
  - [0106] The materials of the coin battery and the battery performances are set forth in Table 2.
  - [Example 14]

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- [0107] In a non-aqueous mixture of EC/PC/DEC=30/5/65 (volume ratio) was dissolved LIPF<sub>6</sub> to give a non-aqueous electrolytic solution of IM concentration. Subsequently, 2.0 wt.% of text-pentylicluses was added to the non-aqueous electrolytic solution.
- [0108] A coin battery was manufactured in the same manner as in Example 11, except for employing the aboveobtained non-aqueous electrolytic solution, and subjected to the evaluation of battery performances.
- [0109] The initial discharge capacity was almost the same as the capacity measured in a battery using an 1M LIPF<sub>6</sub> and EC/PC/DEC (30/6/65, volume ratio) solvent mixture containing no tent-alky/benzene compound [Comparison Example 11].
- [0110] After a 50 cycle charge-discharge procedure, the retention of discharge capacity was 92.1% of the initial discharge capacity (100%). Further, the battery showed good low temperature storage endurance.
  - [0111] The materials of the coin battery and the battery performances are set forth in Table 2.

[Example 15]

- 55 [0112] The procedures of Example 11 were repeated except that (1,1-diethylpropyl)benzene was incorporated in an amount of 2.0 wt.% into the non-aqueous electrolytic solution, to prepare a coin battery.
  - [0113] After the 50 cycle charge-discharge procedure, the retention of discharge capacity was 91,9%.
  - [0114] The materials of the coin battery and the battery performances are set forth in Table 2,

### [Example 16]

- [0115] The procedures of Example 11 were repeated except that a non-aqueous solvent of EC/PC/DEC/DMC (30/50/05, volume ratio) was employed and the artificial graphite was replaced with natural graphite, to prepare a non-aqueous decitorlyle solving and manufacture a coin battery.
- [0116] After the 50 cycle charge-discharge procedure, the retention of discharge capacity was 92.8%.
- [0117] The materials of the coin battery and the battery performances are set forth in Table 2.

### (Example 17)

- [0118] The procedures of Example 11 ware repeated except that a non-aqueous electrolytic solution of 1M LIPFs in ECPPCMEG/DMC (305/50/15, volume ratio) was employed and the LICoC<sub>2</sub> (positive electrode active material) was replaced with LiNe $_2$ CO $_2$ CQ, to manufacture a coin battery.
- [0119] After the 50 cycle charge-discharge procedure, the retention of discharge capacity was 91.1%.
- [0120] The materials of the coin battery and the battery performances are set forth in Table 2.

### (Example 18)

- [0121] The procedures of Example 11 were repeated except that a non-aqueous electrolytic solution of 1M LIBF<sub>x</sub> in ECIPC/DEC/DMC (3005/3005, volume ratio) was employed and the LICoO<sub>2</sub> (positive electrode active material) was replaced with LIMF<sub>x</sub>O<sub>x</sub> to manufacture a coin battery.
  - [0122] After the 50 cycle charge-discharge procedure, the retention of discharge capacity was 92.6%,
  - [0123] The materials of the coin battery and the battery performances are set forth in Table 2.

# 25 [Example 19]

- [0124] The procedures of Example 11 were repeated except that 4-fluoro-tert-penty/benzene was employed as the additive in an amount of 3.0 wt.% to the non-aqueous electrolytic solution, to manufacture a coin battery.
- [0125] After the 50 cycle charge-discharge procedure, the retention of discharge capacity was 92.7%.
- [0126] The materials of the coin battery and the battery performances are set forth in Table 2.

# [Comparison Example 12]

- [0127] The procedures of Example 11 were repeated except that toluene was employed as the additive in an amount of 3.0 wt.% in the non-aqueous electrolytic solution, to manufacture a coin battery.
  - [0128] After the 50 cycle charge-discharge procedure, the retention of discharge capacity was 81.3%.
  - [0129] The materials of the coin battery and the battery performances are set forth in Table 2.

# [Comparison Example 13]

- [0130] The procedures of Comparison Example 11 were repeated except that n-butylbenzene was employed as the additive in an amount of 3.0 wt.% in the non-aqueous electrolytic solution, to manufacture a coin battery.
- [0131] After the 50 cycle charge-discharge procedure, the retention of discharge capacity was 79.7%.
- [0132] The materials of the coin battery and the battery performances are set forth in Table 2.

### [Comparison Example 14]

- [0133] The procedures of Comparison Example 11 were repeated except that di-n-butyl phthalate was employed as the additive in an amount of 3.0 vt.% in the non-equecus electrolytic solution, to manufacture a coin battery.
- [0134] After the 50 cycle charge-discharge procedure, the retention of discharge capacity was 78.1%.
  - [0135] The materials of the coin battery and the battery performances are set forth in Table 2.

### [Comparison Example 15]

- [0136] The procedures of Comparison Example 11 were repeated except that 4-fluorotoluene was employed as the additive in an amount of 3.0 wt.% in the non-aqueous electrolytic solution, to manufacture a coin battery.
  - [0137] After the 50 cycle charge-discharge procedure, the retention of discharge capacity was 80.6%.
  - [0138] The materials of the coin battery and the battery performances are set forth in Table 2.

Table 2

5		Positive electrode Negative electrode	Compound (wt.%)	Electrolytic solution (vol.ratio)	Initial discharge capacity (r.v.) Discharge capacity retention
	Ex.11	LiCoO <sub>2</sub>	tert-pentyl-benzene (2.0)	1M LiPF <sub>6</sub>	1.03
10		Artificial graphite		EC/PC/DEC (30/5/65)	92.8%
	Ex.12	LiCoO <sub>2</sub>	tert-pentyl-benzene (5.0)	1M LIPF <sub>6</sub>	1.02
15		Artificial graphite		EC/PC/DEC (30/5/65)	91.6%
	Ex.13	LICoO <sub>2</sub>	tert-pentyl-benzene (0.5)	1M LIPF <sub>6</sub>	1.01
20		Artificial graphite		EC/PC/DEC (30/5/65)	90.3%
	Com. Ex. 11	LiCoO <sub>2</sub>	None	1M LiPF <sub>6</sub>	1.00
25		Artificial graphite		EC/PC/DEC (30/6/65)	82,6%
	Ex.14	LICoO <sub>2</sub>	1-methyl-4-tert- pentyl-benzene(2.0)	1M LIPF <sub>6</sub>	1.02
30		Artificial graphite		EC/PC/DEC (30/5/65)	92.1%
	Ex.15	LiCoO <sub>2</sub>	(1,1-diethyl-propyl)- benzene(2.0)	1M LIPf <sub>6</sub>	1.02
35		Artificial graphite		EC/PC/DEC (30/5/65)	91.9%
	Ex.16	LiCoO <sub>2</sub>	tert-pentyl-benzene (2.0)	1M LIPF <sub>6</sub>	1.02
40		Natural graphite		EC/PC/DEC/DMC (30/5/30/35)	92.8%
	Ex.17	LiNi <sub>0.8</sub>	tert-pentyl-benzene (2.0)	1M LIPF <sub>6</sub>	1.15
		CO <sub>0.2</sub> O <sub>2</sub>		EC/PC/MEC/DMC	91.1%
45		Artificial graphite		(30/5/50/15)	
	Ex.18	LIMn <sub>2</sub> O <sub>4</sub>	tert-pentyl-benzene (2.0)	1M LiBF <sub>4</sub>	0.99
50		Artificial graphite		EC/PC/DEC/DMC (30/5/30/35)	92.6%
	Ex.19	LiCoO <sub>2</sub>	4-fluoro-tert-pentyl- benzene(3.0)	1M LiPF <sub>6</sub>	1.02
55		Artificial graphite		EC/PC/DEC (30/5/65)	92.7%

#### Table 2 (continued)

	Table 2 (continued)				
5		Positive electrode Negative electrode	Compound (wi.%)	Electrolytic solution (vol.ratio)	Initial discharge capacity (r.v.) Discharge capacity retention
	Com. Ex. 12	LiCoO <sub>2</sub>	toluene (3.0)	1M LIPF <sub>6</sub>	0.98
10		Artificial graphite		EC/PC/DEC (30/5/65)	81.3%
	Com. Ex. 13	LiCoO <sub>2</sub>	n-butyl-benzene (3.0)	1M LIPF <sub>6</sub>	0.97
		Artificial graphite		EC/PC/DEC (30/5/65)	79.7%
15	Com, Ex, 14	LiCoO <sub>2</sub>	di-n-butylphthalate (3.0)	1M LiPF <sub>6</sub>	0.97
		Artificial graphite		EC/PC/DEC (30/5/65)	78.1%
20	Com. Ex. 15	LICoO <sub>2</sub>	4-fluoro-toluene (3.0)	1M LIPF <sub>6</sub>	0.98
		Artificial graphite		EC/PC/DEC (30/5/65)	80.5%

# [Utilization in Industry]

[0139] The present Invention provides a lithium secondary battery showing high safety under overcharge condition, and excellent battery characteristics in cycle property, electric capacity and storage endurance.

# 30 Claims

- A non-equeous electrolytic solution for lithium secondary batteries comprising a non-equeous solvent and an electrolyte, which further contains 0.1 wt.% to 10 wt.% of a tert-alky/benzene compound and 0.1 wt.% to 1.5 wt.% of a biblinery compound.
- The electrolytic solution of claim 1, wherein the tert-alkyribenzene compound is represented by the formula of (R1)
  (IPS) (R9)C-6<sup>1</sup> in which each of R1, R2, and R3 independently is an alkyl group of 1 to 4 carbon atoms, and 41 represents a benzene ring that may have 1 to 5 substitutes to the rinc.
- The electrolytic solution of claim 2, wherein the tert-alkytibenzene compound has no substituent on the benzene ring.
  - 4. The electrolytic solution of claim 2, wherein the tent-alkylbenzene compound is tert-butylbenzene.
- The electrolytic solution of claim 2, wherein the tert-alkylbenzene compound is tert-pentylbenzene.
  - The electrolytic solution of claim 2, wherein the tert-alkylibenzene compound has 1 to 5 hydrocarbyl groups and/ or helogen atoms on the benzene ring as the substituents.
- The electrolytic solution of claim 1, wherein the biphenyl compound is represented by φ²-φ³ in which each of φ²
  and φ³ independently represents a benzene ring that may have 1 to 5 substituents on the ring.
  - The electrolytic solution of claim 7, wherein the biphonyl compound is selected from the group consisting of biphenyl, o-terphenyl, m-terphenyl, p-terphenyl, 4-methylbiphenyl, 4-ethylbiphenyl, and 4-tert-butylbiphenyl.
  - 9. The electrolytic solution of claim 1, wherein the biphenyl compound shows an exidation potential of 4.5 V or lower.

- 10. The electrolytic solution of claim 1, wherein the non-aqueous solvent is at least one compound selected from the group consisting of ethylone carbonate, propylene carbonate, butlyene carbonate, winylene carbonate, repulyroid-actione, dirently estroate, mently estroate, mently estroate, mently estroate, mently estroate, mently estroate, mently estroate, and in a carbonate tertally destroate, 2-retrilyestraphyrotruran, 1,4-dioxane, 1,2-dienthoxyethane, 1,2-dient
- 11. A non-aqueous electrolytic solution for lithium secondary batteries comprising a non-aqueous solvent and an electrolyte, which further contains 0.1 mt.% to 20 mt.% of a test-alky/benzene compound under the condition that the test-alky group has 5 for 3 carbon atoms.
- 12. The electrolytic solution of claim 11, wherein the tert-alkylbenzene compound is tort-pentylbenzene.
- 13. The electrolytic solution of claim 11, wherein the non-aqueous solvent is at least one compound selected from the group consisting of entrylene carbonate, propriene carbonate, butylene carbonate, vinylene carbonate, polytyolocatione, dimentify carbonate, methyl othyl carbonate, diethyl carbonate, tertahydrotruna, 2-methylerterhydrotruna, 1,4-dioxane, 1,2-dimethoxyethane, 1,2-dilethoxyethane, 12-dibutoxyethane, acetonlitrile, methyl propionate, methyl propion
- 14. A lithium secondary battary comprising a positive electrode comprising a compound oxide containing lithium and cobait, nickel or manganese, a negative electrode comprising lithium metal, lithium alloy, or material capable of receiving and releasing lithium, and a non-aqueous electrolytic solution of claim.
- 15. A lithium secondary battery comprising a positive electrode comprising a compound oxide containing lithium and cobalt, nickel or manganese, a negative electrode comprising lithium metal, lithium alloy, or material capable of receiving and releasing lithium, and a non-aqueous electrotic solution of claim 11.

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INTERNATIONAL SEARCH REPORT

# International application No. PCT/JP02/00518 A. CLASSIFICATION OF SUBJECT MATTER Int.Cl H01M 10/40 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) Int.Cl. HOLM 10/40 Documentation saturabed other than minimum documentation to the extent that such documents are included in the fields searched 124cayyo Shinana Roho 1976-1996 Rokai Jilsuyo Shinan Roho 1971-2002 Rokai Jilsuyo Shinan Toroku Koth 1995-2002 Electronic data base consulted during the international search (name of data base and, where oracticable, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Category\* Relevant to claim No. JP 10-275632 A (Hitachi Maxell, Ltd.), 13 October, 1998 (13.10.1998), Par. No. 0012, etc. 1-15 (Family: none) JP 5-36439 A (Sony Corporation), 12 February, 1993 (12.02.1993), Claim 1, etc. (Family: none) A 7-15 JP 2001-167791 A (Ube Industries, Ltd.), P,A 1-15 22 June, 2001 (22.06.2001), Claims 1, 2 (Family: none) JP 2000-058116 A (Sanyo Blectric Co., Ltd.), 25 February, 2000 (25.02.2000), Claims 1 to 4, etc. (Family: none) A 1-15 P,A JP 2001-357876 A (Nippon Steel Chemical Co., Ltd.), 26 December, 2001 (25.12.2001), 1-15 Claim 2, etc. (Family: none)

Further documents are listed in the continuation of Box C. See patent family annex. Special categories of cited documents:

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# INTERNATIONAL SEARCH REPORT

International application No.
PCT/JP02/00518

		PC1/0	1202/00518
	ation). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category	Citation of document, with indication, where appropriate, of the releva	int passages	Relevent to claim No
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